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Application Serial Number: 09/659, 737 Source: Date Processed by STIC:

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.

PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,

2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A

FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 e-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 c-mail help: patin3help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE CHECKER VERSION 3.0 PROGRAM, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings generated for the original version of 37 CFR §§1.821 - 1.825 effective October 1, 1990 (old rules) and the revised version (new rules) effective July 1, 1998 as well as World Intellectual Property Organization (WIPO) Standard ST 25

Checker Version 3.0 replaces the previous DOS-based version of Checker, and is Y2Kcompliant. Checker allows public users to check sequence listings in Computer Readable form (CRF) before submitting them to the United States Patent and Trademark Office (USPTO). Use of Checker prior to filing the sequence listing is expected to result in fewer errored sequence listings, thus saving time and money

Checker Version 3.0 can be down loaded from the USPTO website at the following address: http://www.uspto.gov/web/offices/pac/checker

ERROR DETECTED SUGGESTED CORRECTION

ATTN: NEW PULSE CAL	CTED SUGGESTED CORRECTION	SERIAL NUMBER: 09/659 737
1 Wrapped Nucleics	SES: PLEASE DISREGARD ENGLISH "ALPHA" HEAI	DERS, WHICH WERE INSERTED BY PTO SOFTWARE
	The number/text at the end of each line "wrappe	ed" down to the next line.
	me was retrieved in a wor	rd proposes - B · · · ·
OIPE	Please adjust your right margin to .3, as this will	prevent wrapping
2 Wapped Aminos	The amino acid number/text at the and of analytic	
0 0 0000 A	The amino acid number/text at the end of each li This may occur if your file was retrieved in a wor Please adjust your right margin to 2	ne wrapped down to the next line.
FEB 2 2 2001 o	Please adjust your right margin to .3, as this will	brought have been after creating it.
3 Spreed Line Leng		700
Misaligned Amino A	th The rules require that a line not exceed 72 characters.	cters in length. This includes
Misaligned Amino A		
Numbering		aligned. This may be caused by the use of take
·	between the numbering. It is recommended to del	aligned. This may be caused by the use of tabs lete any tabs and use spacing between the numbers.
5 Non-ASCII	This file was not sayed in a court of	are numbers.
	This file was not saved in ASCII (DOS) text, as re	quired by the Sequence Rules.
_	Please ensure your subsequent submission is sav	red in ASCII text so that it can be processed.
6 Variable Length		
	Sequence(s) contain n's or Xaa's which repri As per the rules, each n or Xaa can only represent	esented more than one residue.
	Please present the maximum number of each resident indicate in the (ix) feature section that	a single residue.
	indicate in the (ix) feature section that some may b	and missing
7 Palentin ver 2.0 "hum		
7 Palentin ver. 2.0 "bug	-3 " · · deficit version 2.0 has caused the <22	20>-<223> section to be miss
	sequence(s) Normally, Patentin w	ould automatically generate this section from the
	previously coded nucleic acid sequence. Please m to the subsequent amino acid sequence. This are	nanually copy the relevant <220 > coor
	to the subsequent amino acid sequence. This app sections for Artificial or Unknown sequences	olies primarily to the mandatory <220> section
•	sections for Artificial or Unknown sequences.	220>~220>
8 Skipped Sequences	Sequence(s) missis a 16 cm.	
(OLD RULES)	Sequence(s) missing. If intentional, please use (2) INFORMATION FOR SEQ ID NO:X:	the following format for each skipped sequence:
	(i) SEQUENCE CHARACTERISTICS (D. T.)	
	(xi) SEQUENCE DESCRIPTION:SEQ ID NO:X:	t any headings under "SEQUENCE CHARACTERISTICS")
	This sequence is intentionally skipped	/
	Please also adjust the "(iii) NUMBER OF SEQUENC	CES: Festioned to include the
9 Skipped Sequences	•	tesponse to include the skipped sequence(s).
(NEW RULES)	Sequence(s) missing. If intentional, please use <210> sequence id number	the following format for each skipped
(337 113223)	<210> sequence id number	supped sequence.
/	<400> sequence id number	
10 Use of n's or Xaa's	Use of n's and/or Yaa's have have	
(NEW RULES)	Use of n's and/or Xaa's have been detected in the Seq Use of <220> to <223> is MANDATORY if n's or Xaa'	quence Listing.
	In <220> to <223> section, please explain toestics of	s are present.
11	In <220> to <223> section, please explain location of r	or Xaa, and which residue n or Xaa represents.
Use of <213>Organism	Sequence(s) are missing this mandatory field	
(NEW RULES)	——— mandatory held	or its response.
12 Use of <220>Feature		• •
(NEW RULES)	Sequence(s) are missing the <220>Feature and a	associated headings
(CONTROLES)	10 12EO 13 MAINDA 1 DRY 11 2013 CODO	AAHOLE: "A
	(See "Federal Register," 6/01/98, Vol. 63,	No. 104 on 20004 on
13 Patentin ver. 2.0 "bug"		
	Please do not use "Copy to Disk" function of Paten file, resulting in missing mandatory numeric identification	illn version 2.0. This causes a corrupted
	file, resulting in missing mandatory numeric identifiers a Instead, please use "File Manager" or any other means to	nd responses (as indicated on raw sequence lighter)
	Instead, please use "File Manager" or any other means t	o copy file to floppy disk.



OIPE

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/659,737

DATE: 02/16/2001 TIME: 11:33:38

Input Set : A:\10489al.app

Output Set: N:\CRF3\02162001\1659737.raw

Does Not Comply Corrected Diskette Needed sce p.6

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3 <110> APPLICANT: Blumenberg, Miroslav
              Gazel, Alix M
      6 <120> TITLE OF INVENTION: GENES AND POLYNUCLEOTIDES ASSOCIATED WITH ULTRAVIOLET
             RADIATION-MEDIATED SKIN DAMAGE AND USES THEREOF
      9 <130> FILE REFERENCE: PC10489A
C--> 11 <140> CURRENT APPLICATION NUMBER: US/09/659,737
C--> 12 <141> CURRENT FILING DATE: 2000-09-11
    14 <150> PRIOR APPLICATION NUMBER: 60/155,029
    15 <151> PRIOR FILING DATE: 1999-09-20
    17 <160> NUMBER OF SEQ ID NOS: 19
    19 <170> SOFTWARE: PatentIn Ver. 2.1
    21 <210> SEQ ID NO: 1
    22 <211> LENGTH: 164
    23 <212> TYPE: DNA
    24 <213> ORGANISM: Homo sapiens
    26 <220> FEATURE:
    27 <221> NAME/KEY: CDS
    28 <222> LOCATION: (2)..(163)
    30 <400> SEQUENCE: 1
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       His Arg Asp Ile Lys Ala Gly Asn Ile Leu Leu Clu Lys Ile Glu
   35 cat gat gac atc tgc aat aaa act ttg aag att aca gat ttt ggg ttg
                                             10
   36 His Asp Asp Ile Cys Asn Lys Thr Leu Lys Ile Thr Asp Phe Gly Leu
                                       25
   39 gcg agg gaa tgg cac agg acc acc aaa atg agc aca gca ggc acc tat
   40 Ala Arg Glu Trp His Arg Thr Thr Lys Met Ser Thr Ala Gly Thr Tyr
                                   40
   43 gcc tgg atg gcc cca gaa g
   44 Ala Trp Met Ala Pro Glu
                                                                         164
   45
          50
   48 <210> SEQ ID NO: 2
   49 <211> LENGTH: 54
   50 <212> TYPE: PRT
  51 <213> ORGANISM: Homo sapiens
  53 <400> SEQUENCE: 2
  54 His Arg Asp Ile Lys Ala Gly Asn Ile Leu Leu Glu Lys Ile Glu
  57 His Asp Asp Ile Cys Asn Lys Thr Leu Lys Ile Thr Asp Phe Gly Leu
                                          10
                                      25
  60 Ala Arg Glu Trp His Arg Thr Thr Lys Met Ser Thr Ala Gly Thr Tyr
                                  40
  63 Ala Trp Met Ala Pro Glu
          50
  68 <210> SEQ ID NO: 3
  69 <211> LENGTH: 145
  70 <212> TYPE: DNA
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RAW SEQUENCE LISTING
PATENT APPLICATION: US/09/659,737

DATE: 02/16/2001
TIME: 11:33:38

Input Set : A:\10489al.app
Output Set: N:\CRF3\02162001\1659737.raw

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   73 <220> FEATURE:
   74 <221> NAME/KEY: CDS
   75 <222> LOCATION: (2)..(145)
   77 <400> SEQUENCE: 3
  78 a\cat egg gac ate aag age gae teg ate etg etg aee eat gat gge agg 49
  79 His Arg Asp Ile Lys Ser Asp Ser Ile Leu Leu Thr His Asp Gly Arg
  82 gtg aag ctg tca gac ttt ggg ttc tgc gcc cag gtg agc aag gaa gtg
                                            1.0
  83 Val Lys Leu Ser Asp Phe Gly Phe Cys Ala Gln Val Ser Lys Glu Val
                                      2.5
  86 ccc cga agg aag tcg ctg gtc ggc acg ccc tac tgg atg gcc cca gag ^{\circ}
  87 Pro Arg Arg Lys Ser Leu Val Gly Thr Pro Tyr Trp Met Ala Pro Glu
                                  40
  91 <210> SEQ ID NO: 4
  92 <211> LENGTH: 48
  93 <212> TYPE: PRT
  94 <213> ORGANISM: Homo sapiens
  96 <400> SEQUENCE: 4
  97 His Arg Asp Ile Lys Ser Asp Ser Ile Leu Leu Thr His Asp Gly Arg
 100 Val Lys Leu Ser Asp Phe Gly Phe Cys Ala Gln Val Ser Lys Glu Val
                                         10
                                      2.5
 103 Pro Arg Arg Lys Ser Leu Val Gly Thr Pro Tyr Trp Met Ala Pro Glu
                                  40
 108 <210> SEQ ID NO: 5
 109 <211> LENGTH: 146
 110 <212> TYPE: DNA
 111 <213> ORGANISM: Homo sapiens
 113 <220> FEATURE:
 114 <221> NAME/KEY: CDS
 115 <222> LOCATION: (2)..(145)
 117 <400> SEQUENCE: 5
 118 t^4 cac agg gac atc aag agt gac tcc atc ctg ctg acc ctc gat ggc agg 49
119 His Arg Asp Ile Lys Ser Asp Ser Ile Leu Leu Thr Leu Asp Gly Arg
122 gtg aag ctc tcg gac ttc gga ttc tgt gct cag atc agc aaa gac gtc
123 Val Lys Leu Ser Asp Phe Gly Phe Cys Ala Gln Ile Ser Lys Asp Val
                                      25
126 cet aag agg aag tee etg gtg gga ace eec tae tgg atg geg eec gag g 146
127 Pro Lys Arg Lys Ser Leu Val Gly Thr Pro Tyr Trp Met Ala Pro Glu
                                 40
131 <210> SEQ ID NO: 6
132 <211> LENGTH: 48
133 <212> TYPE: PRT
134 <213> ORGANISM: Homo sapiens
136 <400> SEQUENCE: 6
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DATE: 02/16/2001

PATENT APPLICATION: US/09/659,737 TIME: 11:33:38 Input Set : A:\10489a1.app Output Set: N:\CRF3\02162001\1659737.raw 140 Val Lys Leu Ser Asp Phe Gly Phe Cys Ala Gln Ile Ser Lys Asp Val 20 25 143 Pro Lys Arg Lys Ser Leu Val Gly Thr Pro Tyr Trp Met Ala Pro Glu 144 35 40 148 <210> SEQ ID NO: 7 149 <211> LENGTH: 3627 150 <212> TYPE: DNA 151 <213> ORGANISM: Homo sapiens 153 <220> FEATURE: 154 <221> NAME/KEY: CDS 155 <222> LOCATION: (868)..(1275) 157 <220> FEATURE: 158 <221> NAME/KEY: CDS 159 <222> LOCATION: (1420)..(1553) 161 <220> FEATURE: 162 <221> NAME/KEY: CDS 163 <222> LOCATION: (1900)..(2026) 165 <220> FEATURE: 166 <221> NAME/KEY: CDS 167 <222> LOCATION: (2105)..(2230) 169 <220> FEATURE: 170 <221> NAME/KEY: CDS 171 <222> LOCATION: (2696)..(2833) 173 <400> SEQUENCE: 7 174 gatetgegae eteetteaga acetgeeaaa atgaetagga aaaatgetgt tteeatagea 60 176 agagecaaaa gagaacatga eggeeetgea eteegggate tetetggeae eagatteeca 120 178 gcccagggga gacacetgaa eceeccagat ggtgacacac etetgtggte etetgtcagg 180 180 gacataacet eccageacag atttgeaaac teeetgetge aggeacaage agggetateg 240 182 ggccccaggt gtggctcccc tgccttggtt cagggagtgg agacacagtt gcccactgct 300 184 ccccacccca etgccaggee tettetgccc ccatgggtcc tggggtgggg gagcettggg 360 186 agttgaagaa tgcctctgac ccagattett caagcagect ctgagetcag aggaagagte 420 188 tgcctcacgg cagcctccct ggggtctagc tgtcaatcgc ccaggaagaa atacccagcg 480 190 cgggaccegg cggggaaget ggcettetet gtetteccag gtgcagcaca gcgagtgtaa 540 192 ggagetgtet tgggeetgee eageetggtg eeetgegggg gaetgetgge acaggaetgt 600 194 gactgggett cagetetgte tgaaaatett tgetteagag caceteeeta gtttgatetg 660 196 ataccecgee tgaccetgee agagteeaga ggteaeggeg geeageeeet geeteeggga 720 198 aggttattcc aaatgeteec acageeetga eeetteetgt tgetttgtee ettgeageee 780 200 aacteetett teegacegee geagaaagae aaceeeceaa geetggtgge caaggeecag 840 202 teettgeeet eggaceagee ggtgggg ace tte age eet etg ace act teg gat 894 203 Thr Phe Ser Pro Leu Thr Thr Ser Asp 204 1 206 acc age age cee cag aag tee ete ege aca gee eeg gee aca gge eag 207 Thr Ser Ser Pro Gln Lys Ser Leu Arg Thr Ala Pro Ala Thr Gly Gln 208 10 15 20 210 ctt cca ggc cgg tct tcc cca gcg gga tcc ccc cgc acc tgg cac gcc 990 211 Leu Pro Gly Arg Ser Ser Pro Ala Gly Ser Pro Arg Thr Trp His Ala 30 35 214 cay ate age ace age aac etg tac etg ecc cag gae eec acg gtt gee

215 Gln Ile Ser Thr Ser Asn Leu Tyr Leu Pro Gln Asp Pro Thr Val Ala

RAW SEQUENCE LISTING

RAW SEQUENCE LISTING PATENT APPLICATION: US/09/659,737

DATE: 02/16/200
TIME: 11:33:38 DATE: 02/16/2001

Input Set : A:\10489al.app

Output Set: N:\CRF3\02162001\1659737.raw

N:\CRF3\02162001\1659737.raw
216
218 dag gat gcc otg gat
218 aag ggt gcc ctg gct ggt gag gac aca ggt gtt gtg aca cat gag cag 1086 219 Lys Gly Ala Leu Ala Gly Glu Asp Thr Gly Val Val Thr His Glu Gln 65
60 and the first
444 LLC dag act acc ++-
222 ttc aag gct gcg ctc agg atg gtg gtg gac cag ggt gac ccc cgg ctg 223 Phe Lys Ala Ala Leu Arg Met Val Val Asp Gln Gly Asp Pro Arg Leu 80
The state of the s
440 CLU CTO GAO See 4
226 ctg ctg gac age tac gtg aag att gge gag gge tee ace gge ate gtc 1182
2 2 2 3 3 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
290 L9C FTG GGG GGG
231 Cys Leu Ala Arg Glu Glu His Ser Gly Arg Gln Val Ala Val Lys Met
110 THE VOL ALA VALUE AND THE
201 9E9 98C CFC 380 100
235 Met Asp Leu Arg Lys Gln Gln Arg Arg Glu Leu Leu Phe Asn Glu 236 125 130
125 125 Pho Acr Ci
230 qtqqqaqac acactanii 135
238 gtgggaggac agggtgggac acacacgggg gcgttgggga tgggcagtga gcagccagcc 1335 240 aggctggaca tctgtgagca ggggcagtgg gtggccatgc gtctgggcac tgtgccagcc 1335 242 actcaggccc ccacctgecc ccag gtg gtg atc atg cgg gac tag gac
242 acteaquee ceacetages agggeagting gtggeeatge gtctgggear tatagetagee 1335
242 acteaggee ceacetgee ceagligt gtg gtg ate atg egg gae tae eag eac 1335 Val Val Ile Met 202 aggest gtg geggee 1335
TTO INCL AID ASD TOP OF THE
440 LLC 330 ata ata
246 ttc aac gtg gtg gag atg tac aag agc tac ctg gtg ggc gag gag ctg 1494 247 Phe Asn Val Val Glu Met Tyr Lys Ser Tyr Leu Val Gly Glu Glu Leu 250 tgg gtg gtg gag atg 155
150 150 TO TO THE TOTAL THE TOTAL TO THE TOTAL THE TOTAL TO THE TOTAL
230 tyg 010 ctc ata 1.
250 tgg gtg ctc atg gag ttc ctg cag gga gga gcc ctc aca gac atc gtc 1542
254 tee caa gte ag gtgggcaget ggggaget 170
254 tcc caa gtc ag gtgggcagct gggagggctg gaccctgagt gcaggctgcc 1593
250 100
258 ctcaccatgg ccctgccagg gcaatgtggt cttctgcctg tggcccagaa gacttgggat 1653 262 gaaggatage ttotaga
260 gcctgggctc ccctgcctgc tggggtaact gagacccagg ggtcttggga gacttgggat 1653 262 gaaggatagc ttctagccaa agctcaggcc ccagttttca ccagggctat gggagaaga 1713
262 gaaggatage ttetageeaa ageteaggee ceagtittea ceaggetat ggeetgaaga 1713 264 tgetgeeaaa eagattgeet gggagetgtg gggeetagea ceaggetat ggeetgaetg 1773 266 cagegageege 2222
and a cradition to the same of
268 gtccag g ctg aat gag gag cag att gcc act gtg tgt gag gct gtg ctg 1893 269 Leu Asn Glu Glu Gln Ile Ala Thr Val Cyc Clu 21 270 Leu Strain Glu
4/4 Cay acc ctr acc t
274 Ala Leu Ala Tyr Leu His Ala Gln Gly Val Tle His Ara atc 1990
270 day dyr. dac too 5+2 -1
277 Lys Ser Asp Ser Ile Leu Leu Thr Leu Asp Gly Arg 2036
270 Sie Sie Map Giv Arg
280 teetgteet ggeacageea egeteeaat teeteetgat ceaceactea eteeetttte 2096 283 Value teetgaa gga teetgaa teetgaa geteegaa teetgaa gga teetga
282 aaccgcag gtg aag ctc tcg gac ttc gga ttc tgt gct cag atc accettttc 2096 283 Val Lys Leu Ser Asp Phe Gly Phe Cys Ala Clarth. agc aaa 2146
Val Lys Leu Ser Asp Phe Gly Phe Cys Ala Gln Ile Ser Lys 286 Gas Grant Care Lys 287 288 288 289 289 289 289 289 289 289 289
286 gac gtc cct aag agg aag tcc ctg gtg gga acc ccc tac tgg atg gct 2194
3 3 3 3 3 4 dec ecc tac tgg atg gct 2194

RAW SEQUENCE LISTING DATE: 02/16/2001 PATENT APPLICATION: US/09/659,737 TIME: 11:33:38

Input Set : A:\10489al.app

Output Set: N:\CRF3\02162001\1659737.raw

```
287 Asp Val Pro Lys Arg Lys Ser Leu Val Gly Thr Pro Tyr Trp Met Ala
                                  245
                                                      250
  290 cet gaa gtg ate tee agg tet ttg tat gee act gag gtaacegtte
  291 Pro Glu Val Ile Ser Arg Ser Leu Tyr Ala Thr Glu
                                                                         2240
                              260
  294 cetecacece ceagacetee caaaageaae ttggcaaetg geagetette tgetgtggee 2300
  296 cctccagtga gctcaccaaa agcageeetg gttttcagag teccaeetag tcaacaceet 2360
  298 teccecttte gatggggetg etettaceca gtgaetttge tgccaggaac gagteetgea 2420
  300 agtgetttee teageteaag ggeagaatgg ggtatggeeg ggeeteetat gtatgatgge 2480
  302 ctttctctga gtgactgaca gctgtgtccc tataggcagt ggtcactcat gcaggcagta 2540
  304 actggccaca gggcaggtga ccaggggagg aaggagacag acccaccaag gagagctggg 2600
  306 geoagetgte eccectecae cactgetgee accagaacge agetaccaat gggecagggt 2660
  308 etggecatgg ggtcagggae atttteetee tgeag 'gtg gat ate tgg tet etg
                                             Val Asp Ile Trp Ser Leu
 312 ggg atc atg gtg att gag atg gta gat ggg gag cca ccg tac ttc agt
 313 Gly Ile Met Val Ile Glu Met Val Asp Gly Glu Pro Pro Tyr Phe Ser
                                                                        2761
                275
                                     280
 316 gas too coa gtg caa gee atg aag agg etc egg gas age eec eea eec
 317 Asp Ser Pro Val Gln Ala Met Lys Arg Leu Arg Asp Ser Pro Pro Pro
             290
                                295
 320 aag ctg aaa aac tot cac aag gte agttggcaca caagggtgeg acetegeaga
 321 Lys Leu Lys Asn Ser His Lys Val
         305
                             310
 324 coccatteet cetgaggeaa ggggaceaga acetgggete ceageatete cettecaetg 2923
 326 aagccacagg gtotgggoto otggaaaagg otootottto occacacaaa accogcacct 2983
 328 gggtgtggag cegeatetae geacaagtte geatgtgege teegacaagt egeeteecae 3043
 330 ggctgtggca ggagagttgc tgcttggcag aagggttgct gcttggcagg cactggtcgg 3103
 332 aagcccagtg gggcccatga gcagggaaag ccaggacacc agcaactccc tgctgtccag 3163
 334 ggagggatcc ggagaagctt cactgagcac aaaccettca accegtgtcg ggagatccat 3223
 336 accatgatte gatgteeetg tecateaegg egagtegget catgeteeat tegttgcaca 3283
 338 coccgacaca gotaagocac agogttocco ttaaagocag tataagtgca tggaagtggt 3343
340 atacatgtaa ccctttttgc caaatcggcc ccaaccccgc aggccttact gtggacgccc 3403
342 cetgetggca ggtcagcacg gggctgataa gtggcaccgc catetggtgg ccaaaacaag 3463
344 aaatgtetea gagggetgaa geeteteete taaaatagea aaaaaacaag agttetgtgg 3523
346 coccaacaca aagetggatg ggaggaccaa caggaaacat ettecaagae aactggteet 3583
348 tggageeege acegetaace ccaaaattag catataaage atge
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352 <211> LENGTH: 311
353 <212> TYPE: PRT
354 <213> ORGANISM: Homo sapiens
356 <400> SEQUENCE: 8
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358 1
                     - 5
                                         10
360 Leu Arg Thr Ala Pro Ala Thr Gly Gln Leu Pro Gly Arg Ser Ser Pro
                                     2.5
363 Ala Gly Ser Pro Arg Thr Trp His Ala Gln Ile Ser Thr Ser Asn Leu
            35
                                 40
366 Tyr Leu Pro Gln Asp Pro Thr Val Ala Lys Gly Ala Leu Ala Gly Glu
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<210> 15 <211> 20 <212> DNA

<213> Homo sapiens

<400> 15

gonacytong ongccatcca

<210> 16 <211> 27 <212> DNA

<213> Homo sapiens

<400> 16 cccgaattca tgcamcanga yathaar

<210> 17 <211> 29 <212> DNA <213> Homo sapiens

<400> 17
cccgaattcg (nacytongg ngccatcca

18

Missing mandatory (2207 to (223) features to explain the "n's" in the sequences. See #10 on the Error Summary Sheet.

Note: Though not shown, seg # 11 has an "n" at position 453, which needs a <220> to <223> explaination.

29

VERIFICATION SUMMARY

DATE: 02/16/2001 PATENT APPLICATION: US/09/659,737 TIME: 11:33:39

Input Set : A:\10489al.app

Output Set: N:\CRF3\02162001\I659737.raw

L:11 M:270 C: Current Application Number differs, Replaced Application Number L:12 M:271 C: Current Filing Date differs, Replaced Current Filing Date L:732 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:11 L:732 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:11 L:732 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:11 L:732 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:11 L:732 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:11 L:1091 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:14L:1091 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:14 L:1091 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:14 L:1091 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:14 L:1091 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:14 L:1100 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:15 L:1100 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:15 L:1100 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:15 L:1100 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:15 L:1100 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:15 L:1109 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:16 L:1109 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:16 L:1109 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:16 L:1109 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:16 L:1109 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:16 L:1118 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:17 L:1118 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:17 L:1118 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:17 L:1118 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:17 L:1118 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:17